

u_{sol} describes the energy of H-bonds to solvent. These three model parameters are obtained by fitting to experimental C_p curves. Best fits on 19 different lysozyme mutants under the same thermodynamic conditions (pH, ionic strength, etc.) reveals that u_{sol} can be treated as constant. Moreover, u_{sol} can be robustly parameterized with as few as five experimental mutants (the standard error of 100 random quintets is <12%). It was observed that a second degree of freedom could be removed due to a linear relationship ($R=0.86$) between the remaining two parameters (δ_{nat} and v_{dha}) indicating that a global balance in enthalpy-entropy compensation must be maintained. Consequently, over a fairly wide range of δ_{nat} values {0.4, 1.6}, the correlation between the experimental and theoretical T_m 's is nearly constant (ranging from 0.68 to 0.72). Using the best parameter set, T_m can be predicted for new lysozyme mutants. Results on a validation set of an additional 81 lysozyme point mutations will be presented. This work is supported by NIH R01 GM073082.

[1] D.R. Livesay, et al. *FEBS Lett.* 576, 468-476(2004), and D.J. Jacobs and S. Dallakayan, *Biophys. J.* 88, 1-13(2005).

1536-Pos Board B380

Computational Studies of Nucleosome and Chromatin Folding

Guohui Zheng, Wilma K. Olson.

Rutgers University, New Brunswick, NJ, USA.

Understanding the packaging of nucleosomes on DNA, including how various proteins interact with DNA, is important for understanding the dynamics of the cell. With this goal in mind, we have developed a shape-based model to map histone-DNA recognition quantitatively in terms of atomic contacts and DNA deformability. This method reduces the complexity of nucleosome structure from 3D visualization to a 2D mapping. Comparison of 32 available nucleosome crystal structures with this approach shows promise in deciphering the sequence-dependent binding mechanism of nucleosomes on DNA. We have also developed a novel Monte-Carlo method, involving multi-scale dinucleotide and dinucleosome modeling, to simulate the communication of proteins over long stretches of chromatin-compacted DNA. We compare our predictions of chromatin looping with recent experimental measurements of enhancer-promoter interactions, focusing on (i) the role of the histone tails in enhancing chromatin looping and (ii) the internal folding structures of chromatin under different ionic conditions.

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In Silico Examination Of The Influence Of Nucleotide Modifications And Magnesium Ions On tRNA Structure And Dynamics

Christian Blau, Gerrit Groenhof, Helmut Grubmüller.

Max Planck Institut für biophysikalische Chemie, Göttingen, Germany.

In our work the influence of chemical modifications and ions on RNA structure and dynamics has been tested. The effect of nucleotide modifications on E. coli and yeast tRNA in solution has been examined with a molecular dynamics approach. Simulations show a decrease of helical content in RNA secondary structure due to those modifications. In another step magnesium ion binding effects on the same tRNA were looked at by performing simulations with and without ions bound to tRNA. Ions coordinating nucleotides in those simulations show them highly affecting local secondary structure motifs. Thus the simulations performed give new hints on the function of nucleotide modifications and ion binding to RNA.

1538-Pos Board B382

Estimating Orientational Entropies At Protein Interfaces

Stephanus M. Fenger, Helmut Grubmüller.

Max-Planck-Institute for Biophysical Chemistry, Göttingen, Germany.

Entropy effects of the surrounding water layer at the protein interface have been studied for a long time, and their relevance e.g. for protein folding is well recognized. In molecular dynamics simulations entropy estimates for surrounding explicit water molecules are difficult to calculate with established methods such as thermodynamic integration. Here we present a new method to calculate the orientational contribution to the solvent entropy near the protein interface. We exploit the permutation symmetry of the Hamiltonian such that we get trajectories of "localized" water molecules. Orientational correlations are unaffected by this transformation, which therefore enables us to obtain spatially resolved entropy estimates for the protein water shell.

1539-Pos Board B383

Hydration Dynamics As Revealed By The Fluorescence Stokes Shift: The Origin Of Slow Hydration Dynamics And Breakdown Of Linear Response

Tanping Li¹, Ali A. Hassanali¹, Dongping Zhong^{1,2}, Sherwin J. Singer^{1,3}.

¹Biophysics program, The Ohio State University, Columbus, OH, USA,

²Department of Physics, The Ohio State University, Columbus, OH, USA,

³Department of Chemistry, The Ohio State University, Columbus, OH, USA.

Hydration dynamics in the immediate vicinity of a protein, probed by time-dependent fluorescence Stokes shift experiments, is critical to understand its biological function. Protein Stokes shifts typically exhibit biphasic relaxation following photo-excitation: fast relaxation occurs on a time scale of several picoseconds while slower components indicate additional hydration dynamics on a time scale of tens of picoseconds, or longer. Theoretical studies using both linear response and non-equilibrium molecular dynamics (MD) calculation qualitatively reproduce the observed biphasic behavior of time dependent Stokes shift for Trp-7 (W7) in myoglobin. Comparison with constrained MD simulations with protein frozen at the instant of photo-excitation reveals the molecular mechanism of slow hydration process and establishes the critical role of protein flexibility. Coupled protein-water motion is shown to be necessary for the observation of the slow component of hydration dynamics. Qualitatively similar results are found for a series of additional cases, such as monellin and staph. nuclease. We illustrate why tracking the separate contributions to the Stokes shift without constrained MD studies may not yield an accurate interpretation of protein hydration dynamics. Additionally, we examine the extent to which protein fluctuations obey Gaussian statistics and the linear response approximation to the Stokes shift is valid. Equilibrium fluctuations of the ground-excited energy difference, which control the absorption and fluorescence line shapes, in the ground and excited electronic states are not independent of each other. We illustrate how small differences from Gaussian statistics in one electronic state can be a signature of very significant deviations from linear response theory, such as isomerization, in the other electronic state.

1540-Pos Board B384

Effect of Temperature on the Structural and Hydrational Properties of Human Islet Amyloid Polypeptide in Water

Maximilian N. Andrews^{1,2}, Ivan Brovchenko¹, Roland Winter¹.

¹Dortmund University of Technology, Dortmund, Germany, ²International Max Planck Research School in Chemical Biology, Dortmund, Germany.

Structural and hydrational properties of full-length human islet amyloid polypeptide 1-37 (hIAPP) were studied in relation to the hydration water properties in a temperature range from 250 to 450 K by MD computer simulations. At all temperatures studied, hIAPP does not adopt a well-defined conformation. The alpha-helical content and the number of intrapeptide H-bonds of hIAPP decrease with temperature. The distribution of residues showing dihedral angles characteristic of beta-sheets and poly(L-proline) II helices along the peptide chain is close to random, whereas a clear trend towards cooperative "condensation" is seen for residues showing alpha-helical dihedral angles. This cooperativity is suppressed by heating or by introducing the native intramolecular disulfide bond. Intrinsic volumetric properties of hIAPP were estimated by taking into account the difference in the volumetric properties of hydration and bulk water. The temperature dependence of the density of hydration water indicates that the effective hydrophobicity of the hIAPP surface is close to that of carbon-like surfaces. Similarly to the case of the A β (1-42) peptide, the thermal expansion coefficient of hIAPP is negative: upon heating, it continuously decreases from $\sim 3 \cdot 10^{-4}$ to $\sim 2 \cdot 10^{-3} \text{ K}^{-1}$. A spanning H-bonded network of hydration water, which covers hIAPP homogeneously at low temperatures, breaks via a quasi-2D percolation transition, whose midpoint is at about 320 K. Approximately at this temperature, the experimentally measured lag time of hIAPP aggregation drops in a drastic way. We discuss the possible role of the temperature-induced percolation transition of hydration water on the conformational changes and aggregation propensity of amyloidogenic peptides.

1541-Pos Board B385

A Modified Primary Hydration Shell (PHS) Method Allows Up To Two Orders Of Magnitude Time Saving In Molecular Dynamics Simulations: Application To Large Systems And Lipid Bilayers

Mehdi Bagheri Hamaneh, Matthias Buck.

Case Western Reserve University, Cleveland, OH, USA.

A realistic representation of water molecules is essential in molecular dynamics simulation of proteins. However, the standard method of solvating biomolecules, i.e. immersing them in a box of water with periodic boundary conditions, is computationally very expensive. The primary hydration shell (PHS) method, developed more than a decade ago [1], uses only a thin shell of water around the system of interest, and so greatly reduces the computational power needed for simulations [2]. The method, however, was not perfect especially when large proteins are concerned. We have modified the PHS method in several ways to improve its performance when large systems are simulated [3]. The model is applied to several systems with different sizes, and both water and protein behaviors are compared with those obtained from standard simulations with periodic boundary conditions and with experimental data. Specifically, Lipari-Szabo order parameters for the proteins of interest are shown to be in good agreement with those derived from standard simulations and NMR relaxation